Interventions for Left Main Coronary Artery Disease

Sujatha Vipperla

1Department of Cardiology, Indus Hospitals, Affiliated to CSI India, SCAI, Visakhapatnam, Andhra Pradesh, India

Address for correspondence Sujatha Vipperla, MD, DM, FIC, Department of Cardiology, Indus Hospitals, Affiliated to CSI India, SCAI, Visakhapatnam 530002, Andhra Pradesh, India (e-mail: sujasri@hotmail.com).

Introduction

Unprotected left main coronary artery (ULMCA) disease constitutes 5 to 7% of the patients undergoing coronary angiography.1 It is associated with a high mortality as it supplies a large area of the myocardium—approximately 84% of the left ventricle in right dominant system and 100% in a left dominant system.2 In CASS (Coronary Artery Surgery Study) registry, mortality in medically treated patients is as high as 63% at 5 years.3

Coronary artery bypass surgery (CABG) has been the standard management for patients with ULMCA disease. Percutaneous coronary intervention (PCI) has emerged as an attractive alternative to CABGs with the advent of drug-eluting stents (DES), development of antiplatelet pharmacology, and adjunctive imaging techniques.

Atherosclerotic plaque in left main coronary artery (LMCA) is qualitatively different from other segments of the coronary tree, with minimal necrotic core and less thin cap fibroatheroma.4 Seventy percent of significant left main (LM) lesions involve the bifurcation. Intimal atherosclerosis is located in areas of low shear stress along the lateral wall extending distally on the lateral walls of the left anterior descending (LAD) and left circumflex (LCX) arteries. Involvement of flow divider (carina) is minimal or absent. Plaque extending into the proximal LAD, LCX, or both may be seen in 90, 66.4, and 62%, respectively.

Current Guidelines for ULMCA Revascularization

Although CABG is class I (level of evidence B), with growing body of evidence for ULMCA intervention, PCI in those patients with SYNTAX (Synergy between Percutaneous Coronary Intervention with Taxus and Cardiac Surgery) score ≤ 22 has equivalent indication class I (level of evidence B) as per current European practice guidelines.5 Patients with SYNTAX scores (22–32) have a class IIa (level of evidence B) indication whereas CABG is preferred in patients with SYNTAX score ≥ 32. AHA/ACC/SCAI (American Heart Association/American College of Cardiology/Society of Cardiovascular Angiography and Interventions) guidelines assigned class IIa recommendation in patients with favorable anatomy (SYNTAX < 22 and ostial or trunk LM disease) and in patients with clinical characteristics that predict an increased risk of adverse surgical outcomes (e.g., moderate severe chronic obstructive pulmonary disease, disability from previous stroke, or previous cardiac surgery; STS-predicted risk of operative mortality > 2%).6

Current Evidence for ULMCA Revascularization

Registry Data

In the large, multinational, “all-comers” IRIS-MAIN (Interventional Research Incorporation Society–Left MAIN Revascularization) registry,7 over the past two decades increasingly complex patients with worsening risk-factor profiles have been recruited with improvements in concomitant medical therapy. Over time, the proportion of PCI treatment has progressively increased to approximately 61%. Risk-adjusted survival, composite outcomes, and repeat revascularization have significantly improved for PCI over time but have relatively remained stable for CABG narrowing the gap in the treatment effect between PCI and CABG from the BMS (bare metal stent) period to the early DES period and then to the late DES period. The risks of mortality and the composite of death, myocardial infarction (MI), or stroke (9.1 vs. 9.7%; hazard ratio [HR] 0.97; 95% confidence interval [CI]: 0.81–1.17; \( p = 0.75 \)) were equivalent between PCI and CABG, but the risks of repeat revascularization and major adverse cardiac and cerebrovascular events (MACCE) were higher in the PCI group than in the CABG group.

The results of DELTA-2 (Drug-Eluting Stent for Left Main Coronary Artery) registry8—an international, all-comers, multicenter registry—were compared with those from the historical DELTA-1 CABG cohort using propensity score stratification. The primary endpoint was the composite of death, MI, or stroke at the median time of follow-up. The distal LMCA was involved in 84.6% of the lesions. The primary endpoint of death, MI, or cerebrovascular accident was lower in the PCI DELTA-2 group compared with the historical DELTA-1 CABG cohort (10.3% vs. 11.6%; \( p = 0.03 \)) at 17 months of
follow up. A favorable result was observed for PCI with respect to cerebrovascular accident whereas CABG was more advantageous with respect to target vessel revascularization.

**Randomized Trials**

The SYNTAX trial included a subset of 705 patients with LM disease. There was no significant difference in MACCE (death, MI, stroke, and repeat revascularization) at 5 years (36.9% in PCI patients and 31% in CABG patients) (HR 1.23; 95% CI: 0.95–1.59; p = 0.12). Mortality was 12.8 and 14.6% in PCI and CABG patients, respectively (HR 0.88 [0.58, 1.32]; p = 0.53). Stroke was significantly increased in the CABG group and repeat revascularization in the PCI arm. MACCE was similar between arms in patients with low/intermediate SYNTAX scores but significantly increased in PCI patients with high scores (≥ 33).

In the EXCEL (Evaluation of XIENCE versus Coronary Artery Bypass Graft Surgery for Effectiveness of Left Main Revascularization) trial, 1,905 patients—wihch included a subset of 705 patients with LM disease. There was no significant difference in MACCE (death, MI, stroke, and repeat revascularization) at 5 years (36.9% in PCI patients and 31% in CABG patients) (HR 1.23; 95% CI: 0.95–1.59; p = 0.12). Mortality was 12.8 and 14.6% in PCI and CABG patients, respectively (HR 0.88 [0.58, 1.32]; p = 0.53). Stroke was significantly increased in the CABG group and repeat revascularization in the PCI arm. MACCE was similar between arms in patients with low/intermediate SYNTAX scores but significantly increased in PCI patients with high scores (≥ 33).

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In the EXCEL (Evaluation of XIENCE versus Coronary Artery Bypass Graft Surgery for Effectiveness of Left Main Revascularization) trial, 1,905 patients—with ULMCA disease with ≥ 70% diameter stenosis, or ≥ 50% or ≤ 70% with either (1) noninvasive evidence of LM ischemia, (2) intravascular ultrasound-derived minimal lumen area (IVUS-MLA) ≤ 6.0 mm², or (3) fractional flow reserve (FFR) ≤ 0.80, SYNTAX score ≤ 32, and clinical and anatomic eligibility for both PCI and CABG as agreed to by the local heart team—were randomized to PCI with fluoropolymer-based cobalt-chromium everolimus-eluting stents or CABG. The primary endpoint, the composite of death from any cause, stroke, or MI at 3 years occurred in 15.4% of patients who underwent PCI group versus 14.7% of the patients who underwent CABG group meeting the noninferiority criterion. The secondary endpoint event of death, stroke, or MI at 30 days was lower in the PCI group (4.9% in the PCI group vs. 7.9% in the CABG group). The secondary endpoint event of death, stroke, MI, or ischemia-driven revascularization at 3 years occurred in fewer patients in PCI group. CABG patients had fewer primary endpoint events than the PCI group patients between 30 days and 3 years after the procedure. Revascularization rates at 3 years were higher with PCI than with CABG (12.6 vs. 7.5%) whereas the rates of early MI and major adverse events were lower with PCI than with CABG (8.1 vs. 23.0%).

The EXCEL trial enrolled predominantly males (76%). Most patients were clinically at low risk (60% had stable ischemic heart disease and most patients had normal ejection fraction). Thirty percent were diabetic. According to the sites, the SYNTAX score was low (≤ 22) in 60.5% of the patients and intermediate (23–32) in 39.5% of the patients. The core laboratory assigned higher scores, low in 35.8% of the patients, intermediate in 40.0%, and high (≥ 33) in 24.2%, the group that met the exclusion criteria. Most of the patients (80.5%) had distal left main bifurcation or trifurcation disease, and two- or three-vessel coronary artery disease was present in 51.3% of the patients. Average 1.9 lesions were treated per patient. A mean of 2.4 stents with a mean total stent length of 49.1 mm were implanted per patient. Femoral access was used in 73% patients, and IVUS guidance was used in 77% of patients. Off-pump CABG was performed in 29%. Internal mammary artery (IMA) was used in 98.8% with bilateral IMA in 28.8% patients.

NOBLE (Nordic-Baltic-British Left Main Revascularization Study) trial randomized patients with significant LM lesion visually assessed stenosis diameter ≥ 50% or FFR ≤ 0.80 and no more than three additional noncomplex lesions (additional non-LM complex lesions were chronic total occlusions, bifurcation lesions requiring two-stent strategies, calcified or tortuous lesions). A total of 1,201 patients were randomly assigned to PCI or CABG. Kaplan-Meier 5-year estimates of MACCE (death from any cause, nonprocedural MI, repeat revascularization, or stroke) were 29% for PCI and 19% for CABG (HR 1.48; 95% CI: 1.11–1.96) with CABG being significantly better than PCI. Though all-cause mortality was not different, the rates of MI, revascularization, and stroke were higher in PCI patients compared with CABG patients. Repeat revascularization was higher due to de novo lesion and target non-LMCA lesion revascularization. Notably 1-year MACCE was same in both the groups. Surprisingly there was no association with SYNTAX score and MACCE.

Twenty percent of the population was female with 15% diabetics. Eighty percent had stable angina and most had preserved ejection fraction. Eighty-eight percent had distal LM lesion. Elective double stenting was used in 35% with culotte being preferred technique. 11% of PCI cases received a first-generation DES in the LM. Kissing balloon inflation (KBI) was done in 55%, and any ostial circumflex post-dilatation was done in 79%. Complete revascularization was achieved in 92%. IVUS was done pre-PCI in 47% and post-PCI in 74%. CABG was done with the on-pump technique in 84%, with 93% patients receiving left internal mammary artery. The number of grafts per patient was one in 4%, two in 52%, three in 39%, four in 4%, and five in < 1% of patients.

**Conflicting Results of EXCEL and NOBLE**

Both were well-conducted robust trials. Both studies included similar patients with similar SYNTAX scores, and distal LM was treated in the majority. The conflicting results could be due to:

1. Differences between studies in patient assessment, risk profiles, trial process, or procedural characteristics.
2. Differences in the stent used: EXCEL used a thin strut fluropolymer-based cobalt chromium everolimus-eluting stents whereas NOBLE used first-generation sirolimus-eluting Cypher stent (11%) or the biolimus-eluting Biomatrix Flex stent (89%), resulting in a substantial difference in stent thrombosis (0.7 vs. 3%). As a result, the rate of stent thrombosis or symptomatic graft occlusion was much higher after CABG than after PCI (5.4 vs. 0.7%) in EXCEL but similar (4 vs. 3%) in NOBLE.
3. Strategy of stenting: Both the trials used provisional stenting approach. In approximately 75% of patients IVUS was used. Data were not available from EXCEL regarding two-stent strategy. Thirty-five percent needed a two-stent strategy with culotte technique being used in the majority in NOBLE.
4. Differences in primary endpoint used: EXCEL used death, MI, and stroke. Investigators included peri-procedural MI.
The EXCEL trial did not include repeat revascularization. NOBLE trial, on the other hand, used only nonprocedural MI in the endpoint, and repeat revascularization was part of primary composite endpoint. In the NOBLE trial, there was not much difference in target LM artery revascularization but more than twofold increase in de novo lesion revascularization in the PCI group with most patients undergoing repeat PCI (96%).

5. Difference in trial duration: EXCEL reported data at 3 years whereas NOBLE reported data at 5 years. Remarkably in NOBLE, the 5-year mortality was not different in PCI and CABG groups whereas death, stroke, and repeat revascularization were higher in PCI group. In fact, data from EXCEL may look similar to NOBLE if repeat revascularization is included with evaluation done at 5 years.

6. An unexplained higher risk of stroke after PCI in NOBLE, which was more than twice after PCI than CABG.

Overall, 4,594 patients from five trials (EXCEL, NOBLE, SYNTAX, PRECOMBAT, Boudriot et al) were included in the meta-analysis by Nerlekar et al, with 50% of patients in the metanalysis undergoing PCI using DES. The prevalence of isolated ULMCA stenosis ranged from 10 to 29%, with between 55 and 80% of patients having a distal bifurcation ULMCA lesion. The primary endpoint of death, MI, and stroke was similar between PCI and CABG (odds ratio [OR] 0.97; 95% CI: 0.79–1.17; \(p = 0.73\)). As expected, CABG was associated with a significant reduction in the risk of repeat revascularization.

**Preprocedural Assessment and Planning for Left Main Intervention**

The LM is the largest bifurcation and provides blood supply to > 50% of the total myocardial mass. It has many unique features, which demand different technical approaches compared with non-LM bifurcations. These include the following:

1. The SB (side branch) is usually the LCX that most often has a large reference diameter and is angled, making it difficult to access with guidewires. Acute occlusion of the LCX usually results in considerable ischemia and may induce acute ischemic mitral regurgitation. The T-shaped bifurcation angle of the LM may also affect implantation technique, and a highly angulated LCX takeoff may impact prognosis after LM stenting.
   a. The presence of calcification in the LM is a frequent occurrence, given that the patients’ mean age with LM disease is higher.
   b. The LM is the only bifurcation where the proximal MB originates directly from the aorta. Possibility of guidewires to go behind LM stent struts, guide catheter distortion of stent, and stent longitudinal compression can increase complexity.
   c. The proximal reference diameter may reach > 5 mm, which is close to the dilatation limit of many coronary stents.
   d. LM trifurcations are seen in approximately 10% of LM cases and may require specific treatment strategies.

**Indications**

Conventionally ULMCA with ≥ 50% diameter stenosis indicates hemodynamic significance needing intervention. Angiographic assessment of LMCA has inherent limitations due to short vessel segment, lack of reference vessel, eccentricity, foreshortening on angiography, and missed ostial disease due to deep catheter engagement. FFR to assess the functional significance of LMCA lesion is emerging as a complimentary modality, especially in equivocal lesions. There is a poor correlation between angiography and FFR with interobserver concordance of only 52% in one study that also found that 23% of patients had an LMCA stenosis ≤ 50% with FFR < 0.80. A meta-analysis of eight trials detected no significant difference in primary endpoint (all cause death, nonfatal MI, and revascularization) between revascularized and deferred groups. The rate of subsequent revascularization was higher in the deferred group. Whether this was due to LMCA intervention was not reported. As underlying plaque composition rather than hemodynamic significance of a lesion that determines events in LMCA FFR-based approach may leave a group of patients at risk of MI, a hypothesis not proven in meta-analysis. FUTURE (Functional Testing Underlying Coronary REvascularization) trial included LMCA disease in 11% of patients and showed a nonsignificant excess of mortality trend in the FFR group and no clinical benefit of FFR in comparison with angioplasty.

Researchers concluded that in complex, high-risk patients, FFR may not help in treatment decisions and may create a false sense of safety. The European Society guidelines advocate FFR-guided strategy in absence of ischemia testing (class IA). An important limitation of LMCA FFR is potential overestimation due to presence of downstream lesions. The pressure wire should be advanced distally in both epicardial branches to determine both FFR values by using manual pull-back with intravenous (IV) adenosine, and the FFR value in a disease-free daughter vessel should be used. In the presence of downstream lesions, an apparent FFR > 0.85 in LMCA can be reassuring. IVUS provides more reliable data through direct lumen visualization, especially in eccentric lesions and in contrast streaming. MLA (minimal luminal area) of 5.9 mm² and a minimum lumen diameter of 2.8 mm had the highest sensitivity and specificity correlating with FFR < 0.75. Park et al proposed an MLA cutoff of 4.5 mm² to predict an FFR ≤0.80 with 77% sensitivity and 82% specificity in Asian populations. A deferral strategy may be safe in the short- to medium-term if the IVUS-derived LMCA MLA is > 6 mm², a cutoff used in the EXCEL trial (Fig. 1).

**Clinical Factors**

Presentation with acute coronary syndrome in patients with LMCA disease tips the balance in favor of intervention rather than surgery. In DELTA registry that included all comers with ULMCA stenosis treated with first-generation DES or CABGs, 13.7% had acute coronary syndrome. No difference in primary endpoint (death, MI, or stroke) was found between the groups though repeat revascularization was higher in the
PCI group. PCI in patients with ACS and LMCA is proven to be safe despite all comers design and high SYNTAX score (mean SYNTAX 38.0 ± 18.2).

**Anatomic Factors**

The MEDINA classification is an angiographic classification of bifurcation lesion taking into account the plaque distribution into the branches with Medina classes 1, 1, 1; 1, 0, 1; and 0, 1, 1 denoting true bifurcation lesions. SYNTAX score, SYNTAX II, NERS, and EuroSCORE are widely used to describe anatomic and clinical complexity. The SYNTAX score was incorporated into European guidelines. However, in both EXCEL and NOBLE trials, SYNTAX score was not helpful in predicting outcomes. The DEFINITION (Definitions and impact of complEx biFurcation lesions on clinical outcomes after percutaNeous coronary InterventIOOn using drug-eluting stenTs)21 criteria are the only specific risk score for LMCA disease. LMCA lesions are classified as simple if SB diameter stenosis is < 70% and lesion length < 10 mm. A complex LM lesion has SB diameter stenosis > 70% and lesion length > 10 mm or if it satisfies two of the following six minor criteria: (1) moderate to severe calcification; (2) multiple lesions; (3) LAD-LCX bifurcation angle > 70 degrees; (4) main vessel reference vessel diameter < 2.5 mm; (5) thrombus-containing lesion; and (6) main vessel lesion length > 25 mm.

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**Technique of Left Main Stenting**

**Access and Guiding Catheter**

Radial access with large lumen 6F guiding catheters can be used. In case of an aorto-ostial ULMCA lesion, a 6F guide catheter is suitable for most of cases, except when debulking is planned and a larger guiding catheter and a stronger support are needed. For very tight aorto-ostial ULMCA lesions or severe damping of the guiding catheter, it is safer to place a guidewire in the guiding catheter, to be able to wire the LMCA LAD or LMCA LCX and promptly disengage the guiding catheter from the LMCA ostium to avoid catheter damping. It is preferable to use a 7F guiding catheter in distal LM when elective double stenting is planned or when > 3.5-mm balloons are needed for kissing. The advantages of a 7F catheter are better visualization, decreased friction during advancement of the stents, accommodation of all sizes of burr if rotablation is required, and allowing for IVUS to be performed in the presence of multiple guidewires.

**Ostial and Shaft Lesions**

The operator must choose the best angiographic view (AP cranial or LAO-cranial) to visualize the ostium of LM and properly position the stent. The implantation of the stent for ostial LM stenosis must be done with a small protrusion into the aorta. After implantation, it is important to post-dilate the stent with properly sized balloons to facilitate good stent apposition at
the ostium and allow reengagement of the ostium with the catheter (►Fig. 2). The results of LM ostial and shaft lesions are gratifying according to DELTA registry\textsuperscript{22} though first-generation DES were used. At 3 years, the rate of major adverse cardiac events (MACE) was 19.1\% versus 28.5\% for distal LM, and the rate of TLR was 4.5\% as against 12.6\% for distal LM.

**Distal Lesions**

**Wiring**

LAD and LCX should be wired with the most difficult branch wired first. Jailing of the wire in the SB during main vessel stenting is important as it keeps the SB open and is the only marker for rewiring when SB is occluded. The SB wire changes bifurcation angle that facilitates rewiring. Polymer-coated wires were more resistant to retrieval damage and were more efficient in crossing the SB ostium than nonpolymer-coated wires,\textsuperscript{23} but another study reported that jailed polymer-jacketed guidewires were associated with procedural MI.\textsuperscript{24}

**Lesion Preparation**

Lesion is predilated with noncompliant and scoring balloons. Rotational atherectomy is needed for severely calcified LM lesions.

**Stenting Strategy—One versus Two Stents**

Several studies have reported that the provisional one-stent approach for distal LM bifurcation was associated with more favorable outcomes compared with two-stent techniques due to lower risk of death, MI, and target vessel revascularization,\textsuperscript{25-27} including a lower risk of stent thrombosis.\textsuperscript{27} A meta-analysis of seven observational studies involving 2,438 patients with mean follow-up of 32 months showed decreased risk of MACE with provisional versus two-stent strategy (20.4 vs. 32.8\%; OR: 0.51; 95\% CI: 0.35–0.73), and also decreased target vessel/target lesion revascularization (10.1 vs. 24.3\%; OR: 0.35; 95\% CI: 0.25–0.49) favoring provisional stenting as against two-stent strategy.\textsuperscript{28} In a nonrandomized, retrospective study including all consecutive patients with 50\% stenosis of the LM treated with DES in nine European centers between 2002 and 2004, 62.5\% underwent provisional stenting whereas 37.5\% were in the two-stent group. After 10 years, no differences in TLR or MACE were detected.\textsuperscript{29}

**Single-Stent Strategy**

European Bifurcation Club (EBC) 12th consensus\textsuperscript{30} recommends a provisional SB stenting approach for LM treatment in most cases (►Fig. 3). It was the strategy used in EXCEL and NOBLE trials with 35\% patients requiring a two-stent strategy in NOBLE trial. A simple lesion defined as SB stenosis < 70\% and lesion length < 10 mm based on criteria described in the DEFINITION study\textsuperscript{21} favors a provisional approach. Other factors such as a small LCX < 2.5 mm in diameter or diminutive LCX in a right dominant coronary system and a wide angle between LAD and LCX favor provisional approach.

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**Fig. 2** (A) Left main ostial stenosis in 45-year-old woman with unstable angina. (B) Positioning of stent in left main in AP cranial view. (C) Stent deployment. (D) Stent boost. (E) Post-dilatation of stent (F) Final result.
Second-generation DES should be the first choice. The stent platform should be selected keeping in mind the maximum expansion capability of the stent. The stent size should be selected according to the distal reference diameter to avoid carinal shift. Single-stent crossover from LM into the LAD is the most common approach (Fig. 4). If the predominant lesion is in the LCX and LAD ostium is spared, provisional stenting from LM toward the LCX can be done (inverted provisional). While planning to stent toward the LAD or LCX, it is important to remember that stenting toward the tightest vessel rather than the largest vessel has better outcomes and reduces the number of stents used.

Proximal Optimization

Proximal optimization (POT) is performed after stenting by inflating a short balloon just proximal to the carina, to change the tubular stent to a tapered device fitting the LM and distal MB either LAD or LCX, respecting the anatomy of the bifurcation core segment. Care must be taken so that at least 6 to 10 mm of stent length is proximal to the carina (the smallest length of commonly available balloons). Careful positioning of the balloon for POT is crucial and may influence the final result: If too distal, it increases the risk of SB occlusion; if too proximal, it has no effect on the stent strut toward the SB. Ideally, the distal shoulder of the balloon should be positioned...
Interventions for Left Main Coronary Artery Disease

just proximal to the carina while the proximal part is still in
the stent to avoid geographical miss. The main problem is
that the positioning of the distal marker compared with the
distal shoulder varies among the different balloons currently
available. Compliant or noncompliant balloons in a 1:1 rela-
tion to proximal reference diameter of the LM can be used.
POT apposes the stent to LM, reduces ellipticity of stented
segment, and prevents accidental abluminal wiring. POT
allows strut protrusion into the SB with larger strut opening
and minimizes carinal shifting for easier guidewire exchange.

After POT there are three options for the SB (►Fig. 4):
1. Leaving the SB alone if the result is acceptable.
2. If intervention is required to the SB, SB is rewired through
the most distal strut (closest to carina) with either a new
wire or by pulling back the MB wire. The jailed wire should
be withdrawn carefully to avoid deep intubation of the
guiding catheter and consequent damage to the stent. SB is
opened with a short noncompliant balloon, and re-POT is
done to restore stent distortion opposite the SB. (POT-side-
POT [PSP])
3. After wire exchange, KBI is performed with two short,
preferably noncompliant balloons sized according to the
distal diameters. Re-POT is performed (POT-KBI-POT).

The clinical outcomes of recent studies comparing KBI
versus non-KBI failed to show its advantage over non-KBI
treatment in terms of the rate of MACEs, despite lowering SB
diameter stenosis at 6- to 12-month follow-up.12

According to the Korean Coronary Bifurcation Stenting
(COBIS) registry, KBI was associated with a higher MACE rate
due to higher TLR rather than death or MI.33 The COBIS II regis-
try demonstrated the opposite result with reduced MACE and
TLR with KBI. Occlusion of the SB after MV stenting was associ-
ated with more frequent cardiac death and MI, which suggested
the importance of KBI to secure SB patency.34 KBI procedure
can be optimized by final POT to correct proximal malap-
position, proper guidewire cross in the distal cell to optimize SB
strut opening and selecting the appropriately sized balloons for
optimizing the geometry.35 To reduce proximal deformation, a
“modified KBI approach” was recently proposed, using asym-
metric inflation pressures: the SB is first inflated to 12 atm, 
then partly deflated back to 4 atm with simultaneous infla-
tion of the MB balloon at 12 atm. The modified FKBI procedure
reduces elliptical stent deformation and optimizes SB access.35

Conversion to Two-Stent Strategies

SB intervention should be performed in patients who develop
ECG changes or ischemic symptoms after MV stenting attrib-
utable to SB compromise. Treating asymptomatic angiog-
raphic stenosis, though done frequently to prevent future
deterioration, is controversial. Pilot studies reported a consid-
erable discrepancy between angiographic stenosis (50%) and
FFR values.36,37 Incorporating the FFR-guided PCI strategy to
treat the LCX reduces the incidence of unnecessary SB inter-
vention and its attendant complications. However, long-term
clinical trials are needed to establish FFR-guided SB approach
in LM provisional stenting approach. SB stenting can be per-
formed (T, T, and protrusion [TAP] or culotte), with a system-
atic final KBI and a final POT. If wire recrosses through distal
strut, T-stenting is preferred, and if wire crosses through proximal strut, TAP or culotte stenting is preferred (►Fig. 5).

Fig. 4 Provisional stenting approach.

Fig. 5 Provisional stenting conversion to two-stent strategy. ECG, electrocardiogram; POT, proximal optimization treatment; SB, side branch; TAP, T-and-protrusion.
Two-Stent Techniques
A complex LMB lesion according to DEFINITION criteria has SB diameter stenosis > 70%, and lesion length > 10 mm may necessitate a two-stent technique. This corresponds to Medina classification 1, 1, 1 or 1, 0, 1 or 0, 1, 1. Large LCX ≥ 2.5 mm in diameter or diseased left dominant coronary system also may necessitate two-stent techniques.

T-stenting offers good SB scaffolding with angles > 70 degrees but could leave a potential gap at SB ostium or protrusion of SB stent into the MB (in the case of TAP). Variation of the T-stenting, the mini-crush technique was developed involving minimal (usually 1–2 mm) retraction of the SB stent into the MB before crushing with a stent or with a balloon followed by stent. It is relatively simple, leaves the SB open, and provides good coverage of the SB though it leaves multiple layers of strut. However, due to the low success rate of final KBI and suboptimal long-term outcome in the absence of KBI, this procedure has been modified by Chen et al as the double kissing (DK)-crush technique.

Double Kissing Crush Technique
This technique consists of stenting the SB, completely crushing the SB stent with main vessel balloon sized 1:1 to the proximal vessel diameter, proximal SB recross, first KBI, MV stenting, second SB recross and second KBI, followed by final POT (Figs. 6, 7). Another key step in the procedure is the alternative inflation with a noncompliant balloon at high pressure (≥ 16 atm) for the SB before each kissing. First kissing can optimize the distorted SB stent, enlarge the cell of the SB stent, and leave only one layer of struts at the ostial SB, which probably facilitates the second kissing after stenting the MV. FKBI was successfully performed in 100% of cases by DK crush.

Culotte Technique
There are two distinct culotte techniques. The first technique is part of the provisional strategy. When the result is unacceptable in the SB following provisional stenting, when wire recrosses through proximal strut, or when the angle between LAD and LCX is < 70, the SB stent is deployed with the proximal part in the LM followed by POT and KBI. The second strategy begins with wiring of MV and SB. The SB is then ideally stented first, from before the bifurcation in the MV, to beyond the diseased segment of the SB, with a wire jailed in the MV. After POT, the MV is then rewired through a distal strut and the jailed wire is removed. Stent struts are opened with a balloon, and SB wire is removed to prevent metal-to-metal jail and the MV is stented from before the bifurcation to beyond the diseased segment in the MV, according to the diameter of the distal vessel. After a further POT, the SB is rewired, and high-pressure (20 atm) individual noncompliant balloon inflations are made in each vessel at the bifurcation point according to the diameter of the branch vessel to ensure good stent strut separation.

Finally, a lower-pressure KBI is made at the bifurcation. A final POT in stented segment proximal to the bifurcation is optional. The proximal MV is covered by two overlapping stents. It may cause intraprocedural acute closure of the MB after SB stenting, which can be catastrophic during interventions for distal LM disease. The proximal double stent layers can lead to delayed endothelialization and subsequent stent thrombosis. Finally, the distal MB stent at the ostial LAD artery can be underexpanded because of the positioning through the SB stent strut.

EBC TWO trial included patients with true bifurcation lesions in which both the main vessel and SB reference diameters were > 2.5 mm and SB ostial disease was ≥ 5 mm in length. There was no difference in the composite endpoint of death, MI, or target vessel revascularization at 12 months between a provisional approach and a systematic culotte stenting. The inclusion of periprocedural biomarker release in composite primary endpoints of trials on bifurcation lesions might favor provisional stenting, though it does not have an independent prognostic significance. A difference in mortality between the two treatment strategies has never been demonstrated, even for bifurcation lesions with large diameter and extension, as in the EBC TWO trial (2.0 vs. 1.1%; p = 0.59). Similarly, in the Nordic-Baltic Bifurcation Study IV that randomized 450 patients with true bifurcation lesions involving a large SB (≥ 2.75 mm) to provisional stenting or a two-stent strategy (using the culotte technique in 65.5% of cases), the rate of cardiac death and all-cause death at 2 years was similar between the two groups. This lack of mortality difference may be due to lack of impact of bifurcation procedures on prognosis, to the low residual risk of patients enrolled in bifurcation trials, to small sample sizes or to the short duration of follow-up. A recent pooled patient-level data analysis from two large bifurcation trials (Nordic bifurcation study and British bifurcation coronary study) with a 5-year follow-up suggested that a provisional approach is associated with lower all-cause mortality than a systematic dual-stenting technique (3.8 vs. 7%; p = 0.04).

DKCRUSH-III study, a randomized multicenter two-stent study that compared DK crush with culotte stenting for patients with distal LM lesions, showed increased rates of MACE in patients randomized to the culotte group as opposed to the DK crush group (23.7 vs. 8.2%, respectively; p  < 0.001), mainly due to increased MI and target-vascular revascularization. Definite ST rate was 3.4% in the culotte group and 0% in the DK crush group. Complex lesions were associated with a higher rate of MACE at 3 years compared with simple lesions, with an extremely higher rate in the culotte group.

The DKCRUSH-V randomized trial included 482 patients with true distal LM bifurcation lesions (Medina 1,1,1 or 0,1,1) to provisional stenting or DK crush stenting. The primary endpoint was the 1-year composite rate of target lesion failure. A planned DK crush two-stent strategy resulted in a lower rate of TLF at 1 year than a PS strategy; 48.3% of provisional stenting group were transferred into two-stent strategy. One-year TLF was higher in PS group than in DK crush group in complex LM lesions (18.2 vs. 7%; HR 0.27; 95% CI: 0.05–0.54), thus favoring a DK crush technique in complex lesions.
Interventions for Left Main Coronary Artery Disease

Fig. 6 Double-kiss (DK) crush technique. (A) Baseline CAG revealing significant LM bifurcation lesion involving proximal LAD and LCX (Medina 1, 1, 1) (B) Significant calcification in LM, LAD, and LCX and IVUS catheter did not cross. (C) Rotablation of LCX. (D) Rotablation followed by balloon dilatation of LAD. (E) After adequate lesion preparation. (F) Stenting of LM-LCX. (G) Crushing of LCX stent with noncompliant balloon placed in LM-LAD. (H) First kissing balloon inflation after LCX recross. (I) Deployment of stent in LM-LAD. (J) Second kissing balloon inflation after high-pressure sequential dilatation of LAD and LCX. (K) POT in LM with short noncompliant balloon. (L) Result.
**Postprocedure Imaging**

FFR post-PCI is used more often to assess the SB to guide intervention in the SB rather than assess the result of stenting LM. Postprocedure imaging with IVUS is strongly advocated. Imaging can identify stent malapposition, dissections, or significant residual disease, which cannot be detected on conventional angiography. Stent underexpansion is the most important cause of DES failure. Kang et al.\(^4^5\) reported the best IVUS-MSA criteria that predicted angiographic restenosis on a segmental basis included 5.0 mm\(^2\) for the LCX ostium, 6.3 mm\(^2\) for the LAD ostium, 7.2 mm\(^2\) for the polygon of confluence POC, and 8.2 mm\(^2\) for the proximal LM above the POC; 33.8% had under expansion of at least one segment, and angiographic ISR was more frequent in lesions with underexpansion of at least one segment versus lesions with no underexpansion (24.1 vs. 5.4%, \(p < 0.001\)).

Two-year major adverse coronary event-free survival rate was significantly lower in patients with underexpansion of at least one segment versus lesions with no underexpansion. A meta-analysis of 10 studies\(^4^6\) compared IVUS-guided with angiography-guided PCI to determine the effect of IVUS on the mortality in patients with LMCA disease. IVUS-guided PCI was associated with significantly lower risks of all-cause death, cardiac death, target lesion revascularization, and in-stent thrombosis. IVUS-guided PCI in LMCA disease significantly reduced the risks of all-cause death by approximately 40% compared with conventional angiography-guided PCI. Current guidelines recommend IVUS-guided PCI in patients with LMCA disease (class IIa, level of evidence B).\(^5\) The high spatial resolution of OCT (optical coherence tomography) provides exquisite detail of stent apposition, coverage, and proximal or distal SB guidewire crossing. A shallow depth of penetration, which limits assessment of plaque burden, and the need to flush the lumen to achieve a blood-free field are important limitations in using OCT for optimizing LMCA treatment. Unlike IVUS, there are no comparable data for OCT-guided PCI in the LMCA.

**Hemodynamic Support**

Patients with ULM disease with ejection fraction < 35% constitute a high-risk subset needing hemodynamic support. This was addressed in PROTECT II which compared the Impella 2.5 with an intra-aortic balloon pump (IABP) in patients with an
average left ventricular ejection fraction of 24%. The 30-day incidence of MACE was not different for patients with IABP or Impella 2.5 hemodynamic support, though a trend for improved outcome was noted for Impella 2.5 supported patients at 90 days.6 The Impella device is recommended for high-risk LM patients with depressed ejection fractions of < 35%.

Conclusion

PCI for ULM disease using second-generation DES is safe, with clinical outcomes comparable to CABG. The gap between both the techniques has narrowed with the new-generation DES and evolution of techniques of stenting bifurcations. Provisional stenting is the favored approach with two-stent techniques reserved for complex cases. Careful preprocedural planning, systematically performed intervention sticking to the basics of POT, improves outcomes. Imaging further optimizes outcomes.

Conflict of Interest

None.

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